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Description

The invention relates to a semiconductor arrangement having nonvolatile memories.

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Various cells which can be used in the production of are known from the prior semiconductors art. US 4,371,883 describes a cell which has a film made of an organic material between two metal electrodes, the electron acceptor forming a charge transfer complex (CT complex) with one of the electrodes, which comprises silver (Ag). The organic copper (Cu) or material US 4,371,883 for described in is example tetracyanoquinodimethane (TCNQ),

tetracyanonaphthoquinodimethane (TNAP), tetracyanoethylene (TCNE), dichlorodicyanobenzoquinone (DDQ), or the derivatives thereof. Using an electric field, the cell can be switched between two states having different resistances (ON state and OFF state), so that these two states can be assessed for example as "O" or "1".

with US 4,371,883 The cell in accordance significant disadvantages, however, so that such a cell is not appropriate for use in microelectronics. One 25 cell in accordance with disadvantage of the US 4,371,883 consists, inter alia, in the fact that the film thickness deemed necessary lies between 1 and 10 µm. The further disadvantage is that the ratio between the resistances of the ON and OFF states is 30 very low and amounts to only 66, and also that the cell in accordance with construction of the not compatible with the customary US 4,371,883 is constructions in microelectronics. Thus, by way of example, electrodes such as gold, magnesium or chromium 35 are avoided in chipmaking. The crucial disadvantage, cell cannot be used is that the however,

nonvolatile memory cell since such a cell undergoes

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transition from the ON state to the OFF state after the electric field has been switched off (US 4,371,883, column 5, lines 15-17). The transition time is dependent on the film thickness. Further embodiments of such cells are described e.g. in US 4,652,894 or 5,161,149.

The object of the present invention is to provide a semiconductor arrangement having a nonvolatile memory cell which enables a high integration density, is compatible with the customary production methods in microelectronics and has improved properties compared with the memory cells in accordance with the prior art. This object has been achieved by means of the subject matter of patent claim 1.

The advantages of the cell construction according to the invention are reversible switchability, a ratio between ON and OFF resistances of up to 1000 or higher need for nondestructive reading since there is no cell operates rewriting after reading, since the according to a resistive principle, scalability up to an area of 40 nm², nonvolatile information storage, functionality down to film thicknesses of approximately 30 nm, a thermal stability up to 350°C, the functional capability of the cell even at a temperature of up to 200°C, good adhesion of the layers to one another, switchability in the presence of air and moisture, formation of the electrical switchable selective chemical substance directly above the electrode, so that in the presence of an insulator, such as e.g. silicon dioxide, the complex is formed only above the electrode, simple and cost-effective production of the complex, and the suitability of the memory cell for production in a plurality of layers, such as e.g. in the Cu damascene technique.

The semiconductor arrangement having a nonvolatile memory cell according to the invention comprises a substrate, which has two electrodes and an organic material situated in between (identified as material X in the drawings), one electrode forming the compound with the organic material. This "compound" may arise with the formation of covalent or ionic bonds, but also with the formation of charge transfer complexes or of weak bonds such as dipole-dipole interactions, etc.

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Apart from organic materials, it is also possible in particular cases to use inorganic or inorganic-organic materials (likewise as material X) in order to form the abovementioned compound. These materials are, in particular, sulphur, selenium or tellurium either in pure or in bonded form (that is to say organocompounds of sulphur, selenium or tellurium, and, if appropriate, oligomers or polymers). Since organic materials are predominantly used, however, the material is defined as an organic material below. The organic material is preferably selected from the following group:

$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	X ₁ X ₂ R ₁ R ₂ R ₃ R ₅ R ₄ R ₃
NC X ₂ X ₂	NC X ₁ NC X ₂
R ₉ R ₁₁ R ₁₂	R_1 CN R_2 CN
R_1 R_2 CN CH CH CN R_3 R_4	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$
R_1 R_2 CN CH CH CN R_6 R_5 R_4	R_2 R_3 R_4 R_4 R_5 R_7 R_6

NC CN N=S n	NC CN S n
$\begin{array}{c c} R_2 \\ R_1 \\ R_2 \\ Z_1 \\ R_4 \\ CN \\ R_5 \\ \end{array}$	$\begin{array}{c c} R_1 & Z_1 \\ R_1 & Z_2 \\ \hline CN & R_3 \end{array}$
$\begin{array}{c c} R_2 \\ & Z_1 \\ & Y \\ & R_4 \\ & R_5 \end{array}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
NC R ₁ R ₂ R ₂ NC NC	NC Y Z ₂

where R_1 , R_2 , R_3 , R_4 , R_5 , R_6 , R_7 and R_8 , independently of one another, have the following meaning:

H, F, Cl, Br, I (iodine), alkyl, alkenyl, alkynyl, O-alkyl, O-alkenyl, O-alkynyl, S-alkyl, S-alkenyl, S-alkynyl, OH, SH, aryl, heteroaryl, O-aryl, S-aryl, NH-aryl, O-heteroaryl, S-heteroaryl, CN, NO₂, -(CF₂)_n-CF₃, -CF((CF₂)_nCF₃)₂, -Q-(CF₂)_n-CF₃, -CF(CF₃)₂, -C(CF₃)₃ and

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- Z	C≡C()	—Q-CH ₂ -CH=CH ₂	O H C-C CH ₂
O CH ₃ C-C -Q CH ₂	Q C-CH₂ —Q CH=CH₂	0,00	°,c-(
_Q C-CH CH-C	O CH ² CH ₂	°,c—CECH	6 cc≡c()
	— Q — СН=СН- (

the following holds true for n: n = 0 to 10

the following holds true for Q: -O-, -S-

5 R_9 , R_{10} , R_{11} , R_{12} may, independently of one another, be: F, Cl, Br, I, CN, NO_2

 R_{13} , R_{14} , R_{15} , R_{16} , R_{17} may, independently of one another, be:

10 H, F, Cl, Br, I, CN, NO₂

 X_1 and X_2 may, independently of one another, be:

CN	R ₁₃ R ₁₄ R ₁₅ R ₁₇ R ₁₆
R ₁₃ Y R ₁₆ R ₁₄ R ₁₅	R ₁₅ R ₁₄
R ₁₅ R ₁₃ R ₁₃ R ₁₆ R ₁₇	

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the following holds true for Y: O, S, Se The following holds true for Z_1 and Z_2 , independently of one another: CN, NO_2 .

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The substrate may be silicon, germanium, gallium arsenide, gallium nitride; an arbitrary material containing an arbitrary compound of silicon, germanium or gallium; a polymer (that is to say plastic; filled or unfilled; e.g. as molding or film), ceramic, glass or metal. Said substrate may also be an already

processed material and contain one to a plurality of layers comprising contacts, interconnects, insulating layers and further microelectronic components.

5 The substrate is in particular silicon which has already been correspondingly subjected to front end of line (FEOL) processing, that is to say already contains electrical components such as transistors, capacitors, etc. - fabricated using a silicon technology. There is 10 preferably an insulating layer situated between the substrate and the nearest electrode; particularly when the substrate is electrically conductive. However, there may also be a plurality of layers between the substrate and the nearest electrode.

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The substrate may serve only as carrier material or electrical function else fulfill an (evaluation, the last-mentioned case, control). For there contacts between the substrate and electrical electrodes applied to the substrate. Said electrical contacts are for example contact holes (vias) filled with an electrical conductor. However, it is possible for the contacts to be effected from lower layers into the upper layers, through metallizations in the edge regions of the substrate or of the chips.

A preferred device of the invention is the so-called hybrid memory, the substrate being processed using customary front end of the line (FEOL) CMOS silicon technology and the memory layer(s) subsequently being applied thereto. However, the substrate, as mentioned above, is not just restricted to this.

The above-described sandwich structure of the memory cell(s), comprising two electrodes and the intervening organic material or the compound formed, can be applied to the substrate not just once but a number of times in a form stacked one above another. This gives rise to a

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plurality of "planes" for the memory cells, each plane comprising two electrodes and the intervening compound (the electrodes adjoin the two areas of the compound). Of course, there may also be a plurality of cells in one plane (cell array). The different planes can be isolated from one another by means of an insulator. It is also possible for only three electrodes, rather than four electrodes, to be used for two planes lying one above the other; in other words, the "middle" electrode is utilized jointly.

It has surprisingly been ascertained that the cell invention the according to the in semiconductor arrangement can retain the applied state without an applied voltage for a very long time, so that the cell can therefore serve as a nonvolatile memory. able to be shown that the semiconductor arrangement according to the invention having the cell according to invention is still clearly readable functionally capable even after thousands of cycles of the ON/OFF alternation and can even retain the applied state for several months. The electrode facing the substrate (identified hereinafter as bottom electrode) preferably comprises at least two layers, in which case layer which is directly in contact with substrate (identified hereinafter as layer 1 of the bottom electrode) may be titanium (Ti), titanium nitride (TiN), tantalum (Ta), tantalum nitride (TaN), tungsten (W), furthermore TiW, TaW, WN or WCN, and IrO, RuO. SrRuO or any desired combination of these including materials in two or more lavers. in combination with the abovementioned Furthermore, layers or materials, thin layers made of Si, TiNSi, SiN or SiCN may also be present. SiO, SiC, Consequently, layer 1 of the bottom electrode itself may comprise more than one layer.

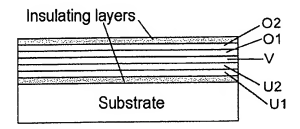
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The abbreviations TiN, TaN, etc. are only symbolic, that is to say that they do not reproduce exact stoichiometric ratios (by way of example, dioxide here is also not identified as SiO2, but as SiO). The ratio of the components can be changed as desired within possible limits. The other (identified hereinafter as layer 2 of the bottom electrode) has a metal, preferably copper, which forms the abovementioned compound with the organic material (material X). This layer (layer 2) which forms the compound may be either pure metal or an alloy of a plurality of metals. What is crucial, however, is that this layer contains a metal which can form the compound with the organic material. The preferred material is copper and its alloys with other metals. Silver or its alloys with other metals is additionally suitable.

Various methods are suitable for depositing abovementioned layers. Said methods may be e.g. PVD, 20 PECVD, vapor deposition, electroplating, electroless plating or atomic layer CVD however, the methods are not just restricted to these methods.

The second electrode (top electrode) may comprise one or a plurality of layers. The second electrode may preferably be aluminum, copper, silver, AlCu, AlSiCu, titanium (Ti), titanium nitride (TiN), tantalum (Ta), tantalum nitride (TaN), tungsten (W), furthermore TiW, TaW, WN or WCN, and IrO, RuO, SrRuO or any desired combination of these materials - including in two or more layers. Furthermore, in combination with the abovementioned layers or materials, thin layers made of Si, TiNSi, SiON, SiO, SiC, SiN or SiCN may also be present. Consequently, layer 1 of the bottom electrode itself may comprise more than one layer.

type of suitable electrodes is not the restricted to the abovementoined materials.



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O2 = layer 2 of the top electrode

01 = layer 1 of the top electrode

V = compound formed

U2 = layer 2 of the bottom electrode

10 U1 = layer 1 of the bottom electrode

The organic material arranged between the electrodes is preferably an electron acceptor, that is to say a molecule having electron-attracting atoms (e.g. -Cl, -F, -Br, -I) or groups (e.g. -CN, -CO-, -NO $_2$) and forms the corresponding compound with the bottom electrode. In particular, those molecules which contain at least one of the abovementioned atoms and/or groups in their skeleton are preferred as electron acceptor. Of course, a plurality of the abovementioned atoms or groups may 20 likewise be present. The preferred organic materials are TCNQ and DDQ. The compound is formed by a selective reaction of the organic material with layer 2 of the bottom electrode, which e.g. contains copper composition of silver. The the bottom 25 contains electrode and of the organic material is not restricted to TCNQ and copper, but rather may on the one hand comprise arbitrary organic materials (which contain at least one of the abovementioned atoms or groups) and on the other hand comprise arbitrary metals. It is only 30 necessary for the electrode to contain a metal which forms the compound with the organic material or with a component of the organic material. The suitable organic material may correspond e.g. to one of the structures

listed in table 1. It is also possible for more than one of the molecules mentioned in table 1 to form the compound with the metal. However, the number of electron acceptors is not limited to the molecules listed in table 1.

Besides the abovementioned electron acceptors, other materials such as e.g. sulphur in elemental form or sulphur-containing organic compounds may also form the compound with the (bottom) electrode (e.g. copper sulphide). Furthermore, by way of example, selenium or selenium-containing compounds or tellurium or tellurium-containing compounds may also form a compound with the bottom electrode.

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The advantageous properties of the cell according to the invention are illustrated in table 2.

Retention time	> 350 days	
Endurance	up to 100.000 cycles	
Threshold voltage (switching)	≥ 2V	
Ratio of the ON and OFF resistances	up to 10.000	
Imprint (writing or erasure)	≥ 1 million pulses	
Read pulses	≥ 10 million pulses	
Scalability	40 nm ²	
Lithographic process directly above the	possible	
cell		

20 Explanations:

Threshold voltage: voltage at which the cell switches from the OFF (ON) to the ON (OFF) state.

Retention time: period of time in which the memory state (ON or OFF) is retained without an applied voltage.

Endurance: number of maximum possible write and erase cycles/pulses.

Imprint: number of maximum possible (single-sided) write or erase pulses without the properties (threshold

voltage, values for ON and OFF resistances, profile of the U-I diagram, etc.) exhibiting a significant, lasting change.

Read: number of maximum possible read pulses.

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A boundary condition for all is that the cells do not break down in the context of the experiments or the electrical values do not exceed specific permitted tolerances.

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The semiconductor arrangement according to the invention may also have a plurality of nonvolatile memory cells and the plurality of cells may be incorporated into the semiconductor arrangement with a high integration density.

The method for producing the semiconductor arrangement is described below.

20 In order to produce the semiconductor arrangement having the memory cell according to the invention, firstly a substrate is provided.

The substrate may be, as described above, silicon, gallium arsenide, 25 germanium, gallium nitride; arbitrary material containing an arbitrary compound of silicon, germanium or gallium; a polymer (that is to say plastic; filled or unfilled; e.g. as molding or film), ceramic, glass or metal. Said substrate may also be an already processed material and contain one to a 30 plurality of layers comprising contacts, interconnects, layers and further microelectronic insulating components.

35 The substrate is in particular silicon which has already been correspondingly subjected to front end of line (FEOL) processing, that is to say already contains electrical components such as transistors, capacitors,

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etc. - fabricated using a silicon technology. There is preferably an insulating layer situated between the substrate and the nearest electrode; particularly when the substrate is electrically conductive. However, there may also be a plurality of layers between the substrate and the nearest electrode.

The substrate may serve only as carrier material or fulfill an electrical function (evaluation, the last-mentioned case, there control). For electrical contacts between the substrate and the electrodes applied to the substrate. Said electrical contacts are for example contact holes (vias) filled with an electrical conductor. However, it is possible for the contacts to be effected from lower layers into the upper layers, through metallizations in the edge regions of the substrate or of the chips.

bottom electrode is applied Firstly, the substrate. There is optionally an insulating layer the situated between the substrate and bottom electrode; in particular, this is a necessity, however, substrate or the topmost layer οf the substrate is electrically conductive. In the case of silicon as substrate, said insulating layer may be e.g. silicon oxide. The bottom electrode introduced into the substrate comprises at least two layers and can be produced by the methods described below.

The electrode may be deposited from the gas phase or from solution. Methods such as e.g. PVD, CVD, PECVD, vapor deposition, electroplating, electroless plating or atomic layer CVD (ALCVD) are suitable for this purpose. The layers U1 and U2 are for example deposited one after the other and subsequently patterned. For this purpose, a photoresist is applied to the layer U2 and this is patterned according to customary methods (exposure, development, etc.). This structure is then

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transferred into the two layers by means of etching using a gas or a gas mixture or else using a liquid or liquid mixture. The etching of the two layers may be effected using the same reagent (gas or liquid) or else may require different reagents.

Apart from patterning by etching, the layers may also be patterned by means of the so-called damascene technique. For this purpose, by way of example, insulating layer (preferably silicon oxide) lying above the substrate is patterned by lithography and etching. After stripping of the photoresist, the two layers are that the trenches or holes deposited, so insulating layer that have arisen during the patterning are completely filled with the electrode materials. That part of these materials which protrudes above the surface of the insulating layer is subsequently ground back. The grinding process may be effected by means of the so-called CMP technique (CMP = chemical mechanical planarization or chemical mechanical polishing). This for interconnects aives rise to, example, contact holes which are filled with the electrode materials and are embedded in the insulating layer or have exactly the same height as the insulating layer.

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Layer 2 of the bottom electrode (U2) is preferably copper or copper-containing and forms the corresponding with the organic material, which is compound subsequently applied. It may also be silver-containing. The organic material may be applied to the electrode (e.g. in a solvent mixture). IF the organic material is solvent mixture comprising at least solvents is preferably used, one of said solvents preferably being acetonitrile or proprionitrile or some other solvent which contains -CN groups. The second solvent is preferably a ketone, an alcohol, an ester, an aromatic, an aliphatic or cycloaliphatic or an ether and mixtures thereof. The following are suitable by way

example: acetone, diethylketone, cyclohexanone, butanone, cyclohexane, gammacyclopentanone, acetate, ethoxyethyl acetate, ethyl butyrolactone, methoxypropyl acetate, ethyoxyethyl proprionate, ethyl alcohol, propyl alcohol, iso-propanol, dibutyl ether, tetrahydrofuran, chlorobenzene, benzyl alcohol. duration of this treatment may be between 10 seconds and 10 minutes. The treatment temperature is between -20 and 100°C, depending on the properties of solvents. Solvent mixtures are also suitable for many 10 substances mentioned in table 1. The proportion of the solvent which contains the -CN group is 0.01 to 65% by volume. Its proportion depends on the composition of the entire solution. This solution may also contain more than two solvents and likewise also more than one 15 organic material (that is to say material X).

Rinsing is then effected using one of the abovementioned solvents, such as acetone for example. This rinsing step serves in particular for removing the excess TCNQ from the substrate, so that only the compound formed remains in the region of the electrode, since the compound can be formed only in this region.

The organic material may also be vapor-deposited onto 25 the bottom electrode. After the vapor deposition, it is necessary to subject the substrate to a treatment in order to produce the compound. It is only after this thermal treatment that the substrate can be rinsed with a solvent in order to remove the excess 30 TCNO. If the organic material is vapor-deposited onto the electrode, it is advantageous if the deposition time is between 2 and 30 min. The pressure to be used lies in a range of between 0.000001 and 200 mbar and the vapor deposition is carried out at a 35 substrate temperature of between -50 and 150°C. It is also possible for not just one but two or more organic materials X to be vapor-deposited onto the electrode simultaneously or one after another.

The properties of the semiconductor arrangement having the memory cell may additionally be improved if the 5 compound formed in the case of a cell produced according to the method described above is subjected to an aftertreatment, to be precise preferably directly after the formation of the compound, sometimes 10 the formation of the compound. during aftertreatment is effected by contacting the compound with a solution of an aftertreatment reagent. particular, amines, amides, ethers, ketones, carboxylic acids, thioethers, esters, aromatics, heteroaromatics, alcohols or various sulphur- or selenium-containing 15 compounds such as e.g. sulphur heterocyclic compounds, compounds with -SO- groups or thiols are appropriate as the aftertreatment reagent, but the number of suitable reagents is not just restricted to these. The reagents may additionally contain unsaturated groups as well as 20 saturated groups. Examples of aftertreatment reagents diethylamine, triethylamine, dimethylaniline, are diphenylamine, dimethylformamide, cyclohexylamine, dimethylacetamide, dimethyl sulfoxide, acetone, 25 diphenylketone, phenyl diethylketone, benzoate, benzofuran, N-methylpyrrolidone, gamma-butyrolactone, toluene, xylene, mesitylene, naphthaline, anthracene, imidazole, oxazole, benzimidazole, benzoxazole, quinoxaline, fulvalene, furan, quinoline, pyrrole, thiophene or diphenyl sulfide. The treatment time is 30 preferably between 15 s and 15 min at a temperature of preferably -30 to 100°C, either in air or under an inert gas, such as e.g. nitrogen or argon.

35 Experience shows that the aftertreatment reagent may be concomitantly incorporated into the memory cell or it may be attached to the cell. The existence of the aftertreatment reagent can be demonstrated for example

after thermodesorption at higher temperatures by means of gas chromatography GC or mass spectroscopy MS. Surprisingly, even very small quantities (from a few of the incorporated or attached aftertreatment reagent can cause significant improvements in properties οf the memory cell. However, the incorporation of the aftertreatment reagent is not a necessity for improving the properties; under certain circumstances, an aftertreatment also suffices for this purpose without an incorporation being detected by means of GC or MS.

As an alternative, the compound may be contacted with gaseous (or vapor) aftertreatment reagent. In air or under an inert gas, such as e.g. nitrogen or argon, the aftertreatment proceeds at a pressure of 0.00001 to 1000 mbar at a substrate temperature of between -30 and 150°C. A thermal step may subsequently follow, but is not always necessary.

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A cell that has been subjected to aftertreatment in this way has an improved (that is to say lower) threshold voltage during switching of the cell by up to 40%, a ratio between the ON and OFF states which is 10 times as high as in the case of a cell that has not been subjected to aftertreatment, and up to 100-fold higher endurance and improved imprint characteristics and an improvement of the layer adhesion by up to 20%.

30 Some of the "aftertreatment reagents" may, however, also be vapor-deposited at the same time as the material X or else directly in succession (they likewise afford the abovementioned advantages), so that they are jointly subjected to the subsequent thermal step.

In a further aspect, the invention relates to an integration concept for a semiconductor arrangement

having a plurality of cells according to the invention. The cell according to the invention may be situated in the semiconductor arrangement between a word line and a bit line which cross one another perpendicularly. The cell is then switched into the ON or OFF state by corresponding voltages being applied to the word line and the bit line. The state of the cell can thus be altered. The ON and OFF states correspond for example to the states having lower and higher electrical resistance, respectively.

In general, the electrodes are produced in such a way that they serve as a word or bit line. However, the situation may also be that an (additional) layer of the top and/or bottom electrode is applied only in the region of the cell - in direct contact with the compound -, that is to say not along the entire interconnect (word or bit line). This relates in particular to the via concept described further below.

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the case of a "crosspoint" construction, individual memory cells lie directly between interconnects that cross one another and form bit and word lines. In order to produce the individual cells, it is possible, by way of example, for the bottom 25 electrodes to be completely covered with the compound and the top electrodes to be applied thereto. crosspoint cells, the size of which is defined solely by the respective widths of the electrodes, thus arise 30 at the crossover points. However, it is also possible for the bottom electrodes not to be covered completely with the compound, but rather only at the locations where the crosspoint cell arises. This becomes possible either by means of the integration method, described later, or by means of a direct patterning of 35 the compound.

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In the case of this crosspoint construction, it is readily possible to provide a plurality of planes of such memory cells in memory cell arrays stacked one above another. Each "plane" of such a memory cell array then contains the associated top and bottom electrodes and also the compound situated in between. It is possible for one electrode to be jointly utilized by two planes, e.g. the top electrode of the first plane may simultaneously serve as the bottom electrode of the overlying second plane. A prerequisite is, of course, that this electrode comprises at least two suitable layers. An insulating layer may also be introduced between two planes, depending on the requirements.

Very high integration densities can thus be achieved, 15 so-called "bit size" being of the order "4F²/n" where magnitude of n is the number individual planes of memory cell arrays that are stacked one above another and "F" denotes the width 20 (smallest possible structure of the technology used).

As an alternative to the abovementioned crosspoint concept, it is possible to produce contact holes directly above the bottom electrode - e.g. in an insulating layer - and then to form the compound in the contact hole directly on the bottom electrode. The size of the cell is then defined by the size of the contact hole (so-called "via concept").

The examples for the integration concept are elucidated in the figures described below, in which:

Figure 1a shows a via concept in which the size of the cell is precisely defined and is not dependent on the size (that is to say width) of the interconnects that cross one another;

- Figure 1b shows an integration concept in which a cell size of approximately $4F^2$ can be achieved (crosspoint concept);
- 5 Figure 1c shows a further integration concept with planes stacked one above another and a bit size of approximately $4F^2/n$; where n = number of planes;
- 10 Figures 2 to 10 show steps which lead to the integration concept in accordance with Figure 1a;
- Figures 11 to 22 show steps which lead to the integration concept in accordance with Figure 1b;
- Figures 23 to 27 show steps which lead to an alternative crosspoint construction, the compound being produced only in the region of the crosspoint cell (and not along an entire electrode as in Figure 14);
- Figures 28 to 44 show a detailed illustration of the method according to the invention.

Figure 2 shows a silicon wafer, in the case of which the FEOL processes have been concluded and the layers depicted there have subsequently been applied. K1 designates a contact (contact hole filled with a conductor material, preferably made of tungsten), B designates layer 1 of the bottom electrode (that is to say U1 according to the previous sketch), C designates a capping layer, I designates an insulating layer and M designates an interconnect. Interconnects M1 or M2 comprise for example layer 1 (= B, e.g. tantalum) and layer 2 (e.g. copper).

K2 designates a contact, that is to say a contact hole which has been filled with the same materials as the interconnect M2. This is effected e.g. in the dual damascene process, in which firstly layer 1 is deposited simultaneously into contact holes K2 and trenches and then layer 2 is deposited. The filled trenches then form the interconnects or electrodes. Layer 1 may also preferably comprise two or more layers (e.g. tantalum nitride and tantalum).

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The capping layer C is preferably Si, TiNSi, SiON, SiO, SiC, SiN, SiCN and any desired combination of these layers or materials.

- 15 D is either a combination of two contacts lying one on top of another or a contact and a pad in order to produce the electrical contact with the substrate and/or with the upper planes.
- 20 Even though the substrate in Figure 2 has been designated as a silicon wafer, the substrate may also be one of the alternatives described earlier. An insulating layer (preferably silicon oxide), is then applied on such a substrate as described in Figure 2.

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Figure 3 shows how the contact holes L are opened in said insulating layer by means of photolithography and etching along the interconnects in order to attain the construction as illustrated in Figure 3a. The capping layer under the contact holes is likewise opened, so that for example the copper surface becomes free there. After the copper surface has been uncovered, the organic material can be applied in order to produce the compound.

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Figure 4 shows how the organic material, TCNQ in the specific case, is deposited onto the substrate surface (Figure 4 relates to vacuum vapor deposition). The

deposition of TCNQ may be effected by means of a vacuum process such as, for example, vapor deposition or by means of a solution of TCNQ. The precise parameters for how the organic material is applied to the electrode are described in the general part of the application. If the organic material from the solvent is brought into contact with the electrode, the desired compound forms selectively only above the electrode. If the organic material is deposited onto the electrode by means of vapor deposition, however, a thermal treatment has to be effected in order to produce the compound.

Figure 5 shows how the compound is formed selectively in the contact holes either after the thermal treatment, if the organic material is vapor-deposited, or directly after the contacting of the electrode with the solution of the organic material. The insulating layer does not react with TCNQ.

Figure 6 shows the substrate surface after rinsing with a solvent, such as acetone for example. The solvent removes the excess organic material that has not formed a compound. For this purpose, the substrate surface may be rinsed by dipping, spraying or spinning (in the spin coater). As a result, the dimensions of the cell are clearly defined and adjacent cells are insulated from one another by the insulating layer.

Figure 7 shows how a further layer of insulating layer can be applied, and in particular how the - newly arisen - substrate surface can be patterned for the production of the interconnects. This may be done using customary lithographic techniques and subsequent etching. The patterning is preferably effected by means of the customary dual CU damascene patterning. In this case, the trenches and contact holes are simultaneously filled with the materials of the corresponding layers and subsequently ground. After the application of the

capping layer, the construction illustrated in Figure 8 is obtained.

The layer B is preferably made of tantalum nitride or is a combination of tantalum and tantalum nitride. The tracks M2 and M3 produced in Figure 8 are perpendicular to one another. The structure illustrated in Figure 1a is thereby obtained (with M2 as bottom electrode and M3 as top electrode).

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A construction illustrated in Figure 9 is obtained by applying a further layer of the insulating layer and repeating the steps that have been elucidated Figures 3 to 8. The interconnect M3 may serve both as electrode for the bottom cell and as bottom electrode for the top cell. M4 is the top electrode of cell and is perpendicular to м3. construction illustrated in Figure 9 is similar Figure 1c, with the difference that Figure 1c shows a stack (construction having more than one cell plane) on the basis of the crosspoint concept and Figure 9 shows a stack on the basis of the via concept. The advantage of the latter construction is that the cell size is precisely defined and that the lateral insulation of the individual memory cells by the insulating layer prevents crosstalk ο£ the adjacent cells. disadvantage of this construction, however, is that the than $4F^2/n$ (lower integration bit size is more density).

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Figure 10 shows how further processing would have to be effected in order to provide an insulating layer between the first and second cell planes (that is to say that M3 would then no longer serve as a common electrode for two cells). After the application of the capping layer to the substrate in accordance with Figure 10, processing would be effected according to Figures 3 to 8 in order to produce the next cell plane.

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Figures 11 to 19 show an integration concept for the semiconductor arrangement according to the present invention, the integration concept enabling a bit size of $4F^2/n$.

Figure 11 shows a substrate similar to that in Figure 2. Figure 2 and Figure 11 make it clear that the substrate may be different. It is also possible to start with a substrate as illustrated in Figure 2. Figure 11a shows the plan view of the structure illustrated in Figure 11.

As already described in Figure 2, the substrate may be either a silicon wafer or silicon, germanium, gallium arsenide, gallium nitride; an arbitrary material containing an arbitrary compound of silicon, germanium or gallium; a polymer, ceramic, glass or metal.

- 20 As illustrated in Figure 12, the capping layer C is opened by means of photolithography and etching in order to uncover the interconnects. The compound is intended to be formed later above said interconnects.
- 25 Figure 13 shows the construction after the deposition of the organic material X. The compound has not yet since formed above the interconnect the material has been vapor-deposited by means of a vacuum method. It is only after the substrate obtained in this way has been subjected to a thermal treatment that the 30 above the interconnect. Since forms compound compound between the metal, e.a. copper, and organic material is formed selectively only above the metal (Figure 14), the opening in the capping layer may be larger than the width of the interconnect M1, the 35 intention being also to take account of the overlay tolerances during the photolithography. The organic material may, as described above, be applied either by

means of a vacuum process or by treatment with a solvent. If the organic material in the solvent is applied to the substrate, the structure depicted in Figure 13 is obviated.

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The substrate is then rinsed with acetone, for example, in order to remove the excess organic material. The result of this step is described in Figure 15. The trapezoidal structure of the compound is only schematic. Once the compound has formed over the entire length of the interconnect, a layer of insulation is applied and ground, e.g. by means of CMP, in order to attain the construction depicted in Figure 16a.

Contact holes for the contacts and also trenches for 15 the interconnects can then be opened according Figure 16b using customary lithography and etching techniques. The interconnects which are now to transversely with respect the formed run Figure 11. 20 depicted as м1 in The interconnects patterning may be effected for example by means of dual Cu damascene patterning. In Figure 16b, T1 is either a contact hole or a trench for a pad and L is a contact hole. T2 is a trench for an interconnect which must exhibit an expansion at least by the magnitude of the 25 alignment tolerances above the contact hole. Figure 16c shows the plan view of the structure illustrated in Figure 16b. The hatched region shows the area where the compound formed becomes visible through the trench T2 30 produced.

As shown in Figure 17, the trenches and holes can be filled and planarized by means of the dual Cu damascene technique. Here B is layer 1 of the top electrode, which preferably comprises tantalum nitride or a combination of tantalum and tantalum nitride. Copper preferably forms layer 2 of the top electrode. In Figure 17, the tracks M1 and M2 are perpendicular to

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one another. Consequently, the memory cells are defined everywhere at the locations where the tracks cross one another. D is either a combination of two contacts K or of a contact and a pad, and serves for wiring the different interconnects in different planes with the substrate.

The construction illustrated in Figure 18 is obtained by applying a further capping layer and subsequently repeating the steps illustrated in Figures 12 to 17. In this Figure 8, the interconnect M2 (comprising for example Ta and Cu or Ta, TaN and Cu) may serve both as top electrode for the bottom cell and as bottom electrode for the top cell. M3 is the top electrode of the top cell and is perpendicular to M2. The construction illustrated in Figure 18 corresponds to Figure 1c.

As shown in Figure 19, an interconnect, such as M2 for example, need not necessarily serve as an electrode for and bottom cells. It is also possible for compound not to be formed on the interconnect M2, but rather for a capping layer and then an insulating layer to be applied and firstly the interconnect plane M3 to contact-connected. and After produced application of a further capping layer, the procedure to Figure 12. In may continue according such construction, each interconnect serves either only as top electrode or as bottom electrode, that is to say no common electrodes for two cell planes lying one above another.

The advantage of this concept is that a bit size of $4F^2/n$ can be achieved. The disadvantage, however, is that the organic material is deposited over the entire interconnect, so that the cells are not isolated from one another by a dielectric. This has the effect that the cells are isolated from one another by a dielectric

only in one direction (e.g. x direction), but not in the y direction, that is to say along the interconnect.

embodiment below shows an alternative production of the integration concept in accordance 5 with Figures 11 to 18 and 19. In this embodiment, after the step illustrated in Figure 15, an insulating layer is deposited and ground back to the level of the formed, which produces the compound construction illustrated in Figure 19a. Afterward, the substrate is 10 etched by means of argon plasma, for example, approximately 20 s to 5 min. In this case, the compound significantly more rapidly than etched insulating layer, so that a height difference produced between the compound layer and the insulation, 15 as illustrated in Figure 20. This selective etching may effected wet-chemically, for example also be treating the substrate with a mixture of ammonia and a solvent, such as dimethylformamide, for example. The 20 purpose of this step is to create space for a further protective layer SC, which is deposited onto compound. Said protective layer is firstly deposited over the whole area, as is illustrated in Figure 21a, but after chemical mechanical planarization (CMP) said layer is maintained only above the interconnect M1 or 25 above the compound (Figure 21b). Said layer preferably comprises the same material as the top electrode or layer 1 of the top electrode if the top electrode comprises a plurality of layers. However, it may also 30 comprise one of the other electrode materials already mentioned. A further insulating layer is subsequently applied in order to attain the construction illustrated in Figure 21c.

35 As illustrated in Figure 22, contact holes for the contacts and trenches for the interconnects or pads can be opened by means of customary lithography and etching techniques, such as e.g. the dual damascene technique,

as already described in Figure 16b. After deposition and grinding of the electrode materials, the structure similar to that in Figure 17 is obtained, with the difference that in the present case (Figure 22a) the layer B above the compound is somewhat thicker. For further construction, a capping layer may again be applied and then the procedure may continue according Figure 15 and arrive at a construction illustrated e.g. in Figure 18 or 22b. Ιf the same material as for the layer B is used as the protective layer SC, the layer B is thicker in Figure 22b than in Figure 18. If different materials are used for the SC, two layers are obtained, lavers and illustrated in Figure 22b. The construction depicted in Figure 22b corresponds to the construction of Figure 18 with an additional SC layer.

The integration concept in accordance with Figures 19a to 22b or 22c differs from the method illustrated in Figures 11a to 19 by virtue of the application of the protective layer selectively to the compound. This has the advantage, in particular, that the compound is protected by said protective layer, e.g. during the etching processes.

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embodiment below shows an alternative to the integration concept for. production of an the semiconductor arrangement according to the invention. In this embodiment, an insulating layer is deposited onto the first interconnect, which also represents the electrode for the cell according invention, and only then is the compound formed (that is to say that the step carried out in Figure 16a is effected before the step carried out in Figure 13 or 14). This concept results in a reduction of the process complexity.

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Figure 23, which approximately in The substrate corresponds to Figures 2 and 11a, has applied to it firstly a capping layer C (cap) and then an insulating layer, preferably made of silicon dioxide, in order to attain the construction as illustrated in Figure 24. The trenches for the later interconnects subsequently opened into said insulating layer by means of photolithography and etching, as illustrated The capping layer under the trenches Figure 25. likewise opened, so that the copper surface becomes free at the locations where the (upper) trenches cross the (underlying) copper tracks.

Onto this substrate surface, the organic material is then deposited onto the copper surface either by means of a vacuum process or by treatment with a solution of the organic material, and the compound is formed. If the deposition of the organic material is carried out by means of a vacuum technique, a thermal treatment has to be effected, which may be carried out e.g. on a hot plate or in a furnace, so that the compound is formed selectively above copper, as illustrated in Figure 26, since the insulating layer does not react with the organic material.

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The substrate surface is then rinsed with a solvent, such as acetone for example. This may be effected by dipping, spraying or in the spin coater. Consequently, the dimensions of the cell are clearly defined and adjacent cells are isolated from one another by the insulating layer, as illustrated in Figure 26. In this case, the compound is not formed along the entire interconnects, but rather only locally at the crossover points.

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The trenches are subsequently filled with the electrode material or materials (if the electrode comprises more than one layer). Afterward, grinding may optionally be

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effected. Figures 27a and 27b show the two possibilities, that is with and without grinding (polishing) of the top electrode.

- 5 The construction essentially corresponding to the integration concept illustrated in Figure 1c is obtained by applying a capping layer and subsequently repeating the steps illustrated in Figures 24 to 27.
- 10 The advantage of this integration concept is that an exact definition of the cell dimensions of the memory cells is possible, so that crosstalk between the cells is largely prevented. It is thus possible to achieve an integration concept having the bit size $4F^2/n$.

It should be noted that the individual layers disclosed in the description may comprise a plurality of layers if this is desirable. The structures illustrated in Figures 28 to 36 elucidate in greater detail how the individual layers can be constructed.

Figure 28 shows the substructure in the case of which FEOL and MOL processes are carried out and provided with contacts K1 as termination. The contacts K1 preferably comprise tungsten.

The construction according to Figure 28 is only an alternative which may serve as a substrate for the intended construction with the memory cells according to the invention.

An insulating layer (J1), preferably SiO, is applied to the substrate. If appropriate, the insulating layer J1 may have applied to it additionally a Cu CMP stop layer S1 made of e.g. silicon carbide (SiC) and, for the protection thereof during the lithography process, additionally a further layer J2, which preferably again

comprises SiO. The state after the deposition of the layers J1, S1 and J2 is illustrated in Figure 29.

The layers J1, S1 and J2 are patterned by means of photolithography and RIE (reactive ion etching), as a result of which the contacts K1 are uncovered, as illustrated in Figure 30.

The two-layered bottom electrode is applied by means of a standard Cu damascene process. Firstly, the barrier 10 deposited, which comprises customary В1 is barrier materials or a combination thereof. After the application of the Cu seed layer, copper is deposited by means of an ECD (electrochemical deposition) process under certain circumstances, is subsequently 15 subjected to thermal aftertreatment. This is followed by the chemical mechanical polishing of copper and of the barrier layer, a high selectivity between the copper and the barrier CMP being necessary. The CMP 20 stop layer S1 is necessary in order to ensure a selective barrier CMP process. The CMP process must otherwise be carried out nonselectively. The structure thus obtained is illustrated in Figure 31.

layer of the interconnect (M1) that has 25 generated in this way may have applied to it a copper diffusion barrier S4, preferably made of HDP (high density plasma) Si and N (not shown in Figures 31 and 32, but later in Figure 41). An insulating layer J3, SiO, is then applied. 30 preferably made of appropriate, the dielectric layer may have applied to it a CMP stop layer S2 made of e.g. SiC, and for the protection thereof during the lithography process, a may additionally further protective layer J4 deposited thereon. The protective layer J4 likewise . 35 structure thus obtained comprises SiO. The illustrated in Figure 32.

Trenches are produced in a subsequent step, trenches in this plane are at an angle of 90° with respect to the M1 tracks in the preceding plane. The trenches produced are illustrated in Figure 33. The layers S2 and J3 and, if appropriate J4 are patterned by means of lithography and RIE (reactive ion etching), result of which the M1 tracks are partly uncovered. The organic material is then deposited on the uncovered locations of the M1 tracks by a method as 10 described in the previous embodiments, in order achieve the compound according to the invention. The structure thus produced is illustrated in Figure 34. It corresponds to Figure 26, with the difference that more details of the layers are shown in Figure 34. subsequently continue 15 procedure may e.g. Figure 27a. After the application of the required number of planes according to Figures 24-27a, a final (topmost) interconnect M2 may be constructed, e.g. by means of whole-area deposition of suitable electrode 20 materials. Customary materials such as e.g. Ti/AlCu/TiN may be used as electrode materials in this case. The structure obtained is illustrated in Figure 35. The patterning here is by means of an RIE process.

- 25 As the last layer, a standard passivation layer P (e.g. SiO, SiN, SiON, SiC and arbitrary combinations of these layers) is deposited and the bonding pads are opened. The structure obtained is illustrated in Figure 36.
- 30 The subsequent figures show a variant of the concept described in Figures 11 to 19, a detailed layer construction being shown below.
- An insulating layer J1, preferably made of SiO, substrate. 35 applied the Ιf appropriate, to insulating layer J1 may have deposited onto additionally a Cu CMP stop layer S1, e.g. made of SiC, and, for the protection thereof during the lithography

process, additionally a protective layer J2, preferably again made of SiO. The structure thus obtained corresponds to the arrangement illustrated in Figure 37. The dielectric is patterned in order to attain a construction as illustrated in Figure 38.

The interconnect forming the bottom electrode deposited by means of a standard Cu damascene process. The bottom electrode comprises at least two layers, as 10 described above. In order to produce the interconnect M1, the barrier layer B1 made of customary barrier materials or a combination thereof is deposited. After application of the Cu seed layer, Cu is deposited by means of an ECD (electrochemical deposition) process 15 under certain circumstances, is subsequently and, subjected to thermal aftertreatment. This is followed by the chemical mechanical polishing of the copper layer and the barrier layer, a high selectivity between the copper and barrier CMP being necessary. construction is illustrated in Figure 39. 20

The organic material can then be deposited selectively into the interconnect, as already explained for Figures 13-15. The structure thus obtained is illustrated in 25 Figure 40. The deposition of the organic material may in Figure 13. effected as described Α (high density plasma) SiN may comprising e.g. HDP subsequently be deposited. This layer serves copper diffusion barrier S4. A further insulating layer which preferably comprises SiO, may 30 deposited onto this layer. If appropriate, a CMP stop layer S3 comprising e.g. SiC may be deposited onto the dielectric layer. In order to protect the S3 layer during the lithography process steps, a protective layer J4, preferably also made of SiO, may 35 additionally be deposited. The structure thus obtained is illustrated in Figure 41.

The next step is to generate the trenches for the interconnects for producing the top electrodes. The structure after etching is illustrated in Figure 42. The trenches to be generated are at an angle of 90° with respect to the M1 tracks in the preceding planes.

After application of the required number of planes, the final (topmost) interconnect M2 may be constructed, as illustrated in Figure 43. After the patterning thereof, a passivation layer P is deposited as the last layer in order to attain the construction illustrated in Figure 44. The passivation layer P may be SiO, SiN, SiON or SiC and an arbitrary combination of these layers.

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In the case of the last plane, the interconnect M1 is treated after the CMP process with the organic material arranged thereon, the compound between the organic material and the metal being produced selectively on the copper tracks. A final interconnect M2, serving as an electrode, is constructed by means of a whole-area deposition of suitable electrode materials, as already described in Figure 34.

25 Instead of silicon dioxide, the so-called "low k" material may also be used as the insulating layer I or J. In this case, k denotes the dielectric constant. Insulating layers which permit a higher signal speed owing to the lower k values in comparison with silicon dioxide are involved in this case.

Examples of such materials are: polyimides, polyquinolines, polymers such as polybenzoxazoles, polyimidazoles, polyquinoxalines, 35 aromatic polyethers, polyarylenes including the commercial dielectric SILK, polynorbornenes; copolymers of the materials mentioned; furthermore porous silicon-containing materials, porous organic

materials (porous polymers), porous inorganic-organic materials.

List of reference symbols

- S Substrate
- K Contact
- D Capping layer
- I Insulating layer, which has a plurality of layers
- IJ Individual layers of the insulating layer I
- M Interconnect
- T Trench for an interconnect
- B Bottom layer of the bottom electrode